

1. A peptide comprising a radiometal-binding moiety, wherein said binding moiety comprises the structure:

wherein  $R^1$ ,  $R^2$ , and  $R^3$  independently are selected from the group consisting of H, lower alkyl, substituted lower alkyl, cycloalkyl, substituted cycloalkyl, heterocycloalkyl, aryl, substituted aryl, heteroaryl, substituted heteroaryl, alkaryl, and a protecting group that can be removed under the conditions of peptide synthesis, provided that at least one of  $R^1$ ,  $R^2$ , or  $R^3$  is H,

 $R^4$ ,  $R^5$ ,  $R^6$ ,  $R^7$ ,  $R^8$ ,  $R^9$  and  $R^{10}$  independently are selected from the group consisting of H, lower alkyl, substituted lower alkyl, aryl, and substituted aryl, or  $R^4$  and  $R^6$  together optionally form a direct bond, and  $R^8$  and  $R^9$  together or  $R^7$  and  $R^9$  together may form a cycloalkyl or substituted cycloalkyl ring, and

wherein NR<sup>10</sup> is located at the N-terminus of said peptide, or is located on an amino acid side chain of said peptide.

- 2. A peptide  $acc\phi rding$  to claim 1, wherein  $R^1$  is H.
- 3. A peptide according to claim 1, wherein  $R^3$  is H.
- 4. A peptide according to claim 1, wherein R4 is H.
- 5. A peptide according to claim 1, wherein  $R^4$  and  $R^6$  together form a direct bond.

- 6. A peptide according to claim 5, wherein R<sup>5</sup> is H.
- 7. A peptide according to claim 1, wherein NR<sup>10</sup> is located at the N-terminus of said peptide.
- 8. A peptide according to claim 1, wherein NR<sup>10</sup> is located on an amino acid side chain of said peptide.
- 9. A peptide according to claim 2, wherein  $R^2$  is lower alkyl or substituted or unsubstituted phenyl.
  - 10. A peptide according to claim 9, wherein R<sup>2</sup> is H.
- 11. A peptide according to claim 10, wherein R<sup>3</sup> is
- 12. A peptide according to claim 11, wherein  $\mathbb{R}^4$  and  $\mathbb{R}^6$  together form a direct bond.
- 13. A peptide according to claim 12, wherein R<sup>5</sup> is H.
- 14. A peptide according to claim 13, wherein  $\mathbb{R}^7$ ,  $\mathbb{R}^8$ , and  $\mathbb{R}^9$  each are H.
- 15. A peptide according to claim 14, wherein R<sup>2</sup> is phenyl.
- 16. A peptide according to claim 14, wherein  $\mathbb{R}^2$  is methyl.
- 17. A peptide according to claim 1, wherein  $\mathbb{R}^8$  and  $\mathbb{R}^9$  are methyl.
- 18. A peptide according to claim 1, further comprising a bound metal atom.

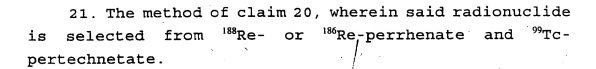
- 19. A peptide according to claim 18, wherein said metal atom is selected from the group consisting of 99mTc, 186Re, and 188Re.
- 20. A method of preparing a metal-chelating composition, comprising contacting a solution of a peptide comprising a radiometal-binding moiety with stannous ions, wherein said binding moiety comprises the structure:

wherein R<sup>1</sup>, R<sup>2</sup>, and R<sup>3</sup> independently are selected from the group consisting of H, lower alkyl, substituted lower alkyl, cycloalkyl, substituted cycloalkyl, heterocycloalkyl, aryl, substituted aryl, heteroaryl, substituted heteroaryl, alkaryl, and a protecting group that can be removed under the conditions of peptide synthesis, provided that at least one of R<sup>1</sup>, R<sup>2</sup>, or R<sup>3</sup> is H,

 $R^4$ ,  $R^5$ ,  $R^6$ ,  $R^7$ ,  $R^8$  and  $R^{10}$  independently are selected from the group consisting of H, lower alkyl, substituted lower alkyl, aryl, and substituted aryl, or  $R^4$  and  $R^6$  together optionally form a direct bond, and  $R^8$  and  $R^9$  together or  $R^7$  and  $R^9$  together may form a cycloalkyl or substituted cycloalkyl ring, and

wherein NR<sup>10</sup>/is located at the N-terminus of said peptide, or is located on an amino acid side chain of said peptide,

and then contacting said solution with a radionuclide and recovering the radiolabeled peptide.



22. A method of imaging a tumor, an infectious lesion, a myocardial infarction, a clot, atherosclerotic plaque, or a normal organ or tissue, comprising administering to a human patient a radiolabeled peptide, together with a pharmaceutically acceptable carrier, and, after a sufficient time for said radiolabeled peptide to localize and for non-target background to clear, the site or sites of accretion of said radiolabeled peptide are detected by an external imaging camera,

wherein said radiolabeled peptide is prepared by contacting a solution of a peptide with stannous ions, wherein said peptide comprises a radiometal-binding moiety comprising the structure:

wherein R<sup>1</sup>, R<sup>2</sup>, and R<sup>3</sup> independently are selected from the group consisting of H, lower alkyl, substituted lower alkyl, cycloalkyl, substituted cycloalkyl, heterocycloalkyl, aryl, substituted aryl, heteroaryl, substituted heteroaryl, alkaryl, and a protecting group that can be removed under the conditions of peptide synthesis, provided that at least one of R<sup>1</sup>, R<sup>2</sup>, or R<sup>3</sup> is H,

 $R^4$ ,  $R^5$ ,  $R^6$ ,  $R^7$ ,  $R^8$ ,  $R^9$  and  $R^{10}$  independently are selected from the group consisting of H, lower alkyl, substituted lower alkyl, aryl, and substituted aryl, or  $R^4$  and  $R^6$  together optionally form a direct bond, and  $R^8$  and  $R^9$  together may form a cycloalkyl or substituted cycloalkyl ring, and

wherein NR<sup>10</sup> is located at the N-terminus of said peptide, or is located on an amino acid side chain of said peptide,

and then contacting said solution with a radionuclide and recovering the radiolabeled peptide.

A peptide according to claim 1, wherein said peptide is selected from the group consisting of: (Chel)  $\gamma$ AbuNleDHF<sub>d</sub>RWK-NH<sub>2</sub>, (Chel) γAbuHSDAVFTDNYTRLRKQMAVKKYLNSILN-NH, KPRRPYTDNYTRLRK (Chel) QMAVKKYLNSILN-NH2, (Chel) γAbuVFTDNYTRLRKQMAVKKYLNSILN-NH2, (Chel)  $\gamma$ AbuYTRLRKQMAVKKYLN\$ILN-NH<sub>2</sub>. HSDAVFTDNYTRLRK (Chel) QMAVKKYLNSILN-NH2, <GHWSYK(Chel)LRPG-NH2, <GHYSLK(Chel)WKPG-NH2, AcNaldCpadWdSRKd(Chel)LRPAJ-NH2, (Chel)  $\gamma$ AbuSYSNleDHF<sub>d</sub>RWK- $\gamma$ H<sub>2</sub>, (Chel)  $\gamma$ AbuNleDHF<sub>d</sub>RWK- $\gamma$ H<sub>2</sub>, (Chel) NleDHF,RWK-NH2 , Ac-HSDAVFTENYTKLRK (Chell) QNleAAK<u>KYLND</u>LKKGGT-NH,, (Chel) γAbuHSDAVFTDNYTRIRKQMAVKKYLNSILN-NH<sub>2</sub>, (Chel) γAbuVFTDNYTRĽRKQMAVKKYLNSILN-NH<sub>2</sub>, (Chel)  $\gamma$ AbuNleDHF<sub>d</sub>RWK-NH<sub>2</sub>°, <GHWSYK (Chel) LRPG-NH<sub>3</sub>,  $\verb|-chel| \verb||| \mathsf{Chel}| \verb||| \mathsf{WKPG-NH}_2, \\ | \mathsf{-chel}| \mathsf{-chel}| \mathsf{WKPG-NH}_2, \\ | \mathsf{-chel}| \mathsf{-c$  $Nal_dCpa_dW_dSRK_d(Chel)WKPG-NH_2$ , < GHWSYK, (Chel) LRPG-NH<sub>2</sub>, AcNaldCpadWdSRKd (Chel/LRPAd-NH2, AcNaldCpadWdSRKd (Chell)LRPAd-NH, AcNal<sub>d</sub>Cpa<sub>d</sub>W<sub>d</sub>SRK<sub>d</sub> (Chel) LRPA<sub>d</sub>-NH<sub>2</sub>, <GHWSYK (Chel) LRPG-NH<sub>2</sub>, Ack(Chel)FdCFWdKTCT-OH, Ack(Chel)DFdCFWdKTCT-OH, Ack(Chel)FdCFWdKTCTfol, Ack(Chel)DFdCFWdKTCT-ol, (Chel) DF<sub>d</sub>CFW<sub>d</sub>KTCT-OH, K(Chel) DF<sub>d</sub>CFW<sub>d</sub>KTCT-ol, K(Chel)KKFdCFWdKTCT-O1, K(Chel)KDFdCFWdKTCT-OH, K(Chel)DSFdCFWdKTCT-OH, K(Chel)DFdCFWdKTCT-OH, K(Chel)DF<sub>d</sub>CFW<sub>d</sub>KTCD-NH<sub>2</sub>, K(Chel)DF<sub>d</sub>CFW<sub>d</sub>KTCT-NH<sub>2</sub>, K(Chel) KDF<sub>d</sub>CFW<sub>d</sub>KTCT-NHNH<sub>2</sub>, Ack(Chel) F<sub>d</sub>CFW<sub>d</sub>KTCT-NHNH<sub>2</sub>, K(Chel)FdCFWdKTCT-ol, and FdCFWdKTCTK(Chel)-NH2, wherein (Chel) is said radiometal-binding moiety.

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